§514.106

publication as described in paragraph (a)(1) of this section.

[40 FR 13825, Mar. 27, 1975, as amended at 51 FR 7392, Mar. 3, 1986; 64 FR 63203, Nov. 19, 1999]

§514.106 Approval of supplemental applications.

- (a) Within 180 days after a supplement to an approved application is filed pursuant to §514.8, the Commissioner shall approve the supplemental application in accordance with procedures set forth in §514.105(a)(1) and (2) if he/she determines that the application satisfies the requirements of applicable statutory provisions and regulations
- (b) The Commissioner will assign a supplemental application to its proper category to ensure processing of the application.
- (1) Category I. Supplements that ordinarily do not require a reevaluation of any of the safety or effectiveness data in the parent application. Category I supplements include the following:
- (i) A corporate change that alters the identity or address of the sponsor of the new animal drug application (NADA).
- (ii) The sale, purchase, or construction of manufacturing facilities.
- (iii) The sale or purchase of an NADA.
- (iv) A change in container, container style, shape, size, or components.
- (v) A change in approved labeling (color, style, format, addition, deletion, or revision of certain statements, e.g., trade name, storage, expiration dates, etc).
- (vi) A change in promotional material for a prescription drug not exempted by §514.8(a)(3)(i) and (a)(3)(ii).
- (vii) Changes in manufacturing processes that do not alter the method of manufacture or change the final dosage form.
- (viii) A change in bulk drug shipments.
- (ix) A change in an analytical method or control procedures that do not alter the approved standards.
- (x) A change in an expiration date.
- (xi) Addition of an alternate manufacturer, repackager, or relabeler of the drug product.

- (xii) Addition of an alternate supplier of the new drug substance.
- (xiii) A change permitted in advance of approval as listed in §514.8(d).
- (xiv) Changes not requiring prior approval which are listed under §514.8(a)(5) when submitted as supplemental applications.
- (2) Category II. Supplements that may require a reevaluation of certain safety or effectiveness data in the parent application. Category II supplements include the following:
- (i) A change in the active ingredient concentration or composition of the final product.
- (ii) A change in quality, purity, strength, and identity specifications of the active or inactive ingredients.
- (iii) A change in dose (amount of drug administered per dose).
- (iv) A change in the treatment regimen (schedule of dosing).
- (v) Addition of a new therapeutic claim to the approved uses of the product.
- (vi) Addition of a new or revised animal production claim.
 - (vii) Addition of a new species.
- (viii) A change in the prescription or over-the-counter status of a drug product.
- (ix) A change in statements regarding side effects, warnings, precautions, and contraindications, except the addition of approved statements to container, package, and promotional labeling, and prescription drug advertising.
- (x) A change in the drug withdrawal period prior to slaughter or in the milk discard time.
- (xi) A change in the tolerance for drug residues.
- (xii) A change in analytical methods for drug residues.
- (xiii) A revised method of synthesis or fermentation of the new drug substance.
- (xiv) Updating or changes in the manufacturing process of the new drug substance and/or final dosage form (other than a change in equipment that does not alter the method of manufacture of a new animal drug, or a change from one commercial batch size to another without any change in manufacturing procedure), or changes in the methods, facilities, or controls used for

Food and Drug Administration, HHS

the manufacture, processing, packaging, or holding of the new animal drug (other than use of an establishment not covered by the approval that is in effect) that give increased assurance that the drug will have the characteristics of identity, strength, quality, and purity which it purports or is represented to possess.

[55 FR 46052, Nov. 1, 1990; 55 FR 49973, Dec. 3, 1990; 56 FR 12422, Mar. 25, 1991]

§514.110 Reasons for refusing to file applications.

- (a) The date of receipt of an application for a new animal drug shall be the date on which the application shall be deemed to be filed.
- (b) An application for a new animal drug shall not be considered acceptable for filing for any of the following reasons:
- (1) It does not contain complete and accurate English translations of any pertinent part in a foreign language.
- (2) Fewer than three copies are submitted.
- (3) It is incomplete on its face in that it is not properly organized and indexed.
- (4) On its face the information concerning required matter is so inadequate that the application is clearly not approvable.
- (5) The new animal drug is to be manufactured, prepared, propagated, compounded, or processed in whole or in part in any State in an establishment that has not been registered or exempted from registration under the provisions of section 510 of the act.
- (6) The sponsor does not reside or maintain a place of business within the United States and the application has not been countersigned by an attorney, agent, or other representative of the applicant, which representative resides in the United States and has been duly authorized to act on behalf of the applicant and to receive communications on all matters pertaining to the application.
- (7) The new animal drug is a drug subject to licensing under the animal virus, serum, and toxin law of March 4, 1913 (37 Stat. 832; 21 U.S.C. 151 *et seq.*). Such applications will be referred to the U.S. Department of Agriculture for action

- (8) It fails to include, with respect to each nonclinical laboratory study contained in the application, either a statement that the study was conducted in compliance with the good laboratory practice regulations set forth in part 58 of this chapter, or, if the study was not conducted in compliance with such regulations, a brief statement of the reasons for the noncompliance.
 - (9) [Reserved]
- (10) The applicant fails to submit a complete environmental assessment under §25.40 of this chapter or fails to provide sufficient information to establish that the requested action is subject to categorical exclusion under §25.30 or §25.33 of this chapter.
- (c) If an application is determined not to be acceptable for filing, the applicant shall be notified within 30 days of receipt of the application and shall be given the reasons therefore.
- (d) If the applicant disputes the findings that his application is not acceptable for filing, he may make written request that the application be filed over protest, in which case it will be filed as of the day originally received.

[40 FR 13825, Mar. 27, 1975, as amended at 50 FR 7517, Feb. 22, 1985; 50 FR 16668, Apr. 26, 1985; 62 FR 40600, July 29, 1997]

§514.111 Refusal to approve an application.

- (a) The Commissioner shall, within 180 days after the filing of the application, inform the applicant in writing of his intention to issue a notice of opportunity for a hearing on a proposal to refuse to approve the application, if the Commissioner determines upon the basis of the application, or upon the basis of other information before him with respect to a new animal drug, that:
- (1) The reports of investigations required to be submitted pursuant to section 512(b) of the act do not include adequate tests by all methods reasonably applicable to show whether or not such drug is safe for use under the conditions prescribed, recommended, or suggested in the proposed labeling thereof; or
- (2) The results of such tests show that such drug is unsafe for use under such conditions or do not show that